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- CMP ABRASIVE, LIQUID ADDITIVE FOR CMP ABRASIVE AND METHOD FOR POLISHING (54)SUBSTRATE
- A CMP abrasive comprising a cerium oxide slurry containing cerium oxide particles, a dispersant and water, and a liquid additive containing a dispersant and water; and a liquid additive for the CMP abrasive. A method for polishing a substrate which comprises holding a substrate having, formed thereon, a film to be pollshed against a polishing pad of a pollshing platen, followed by pressing, and moving the substrate and the

pollshing platen while supplying the above CMP abrasive in between the film to be polished and the polishing pad to thereby polish the film to be polished. The CMP abrasive and the method for polishing can be used for polishing a surface to be polished such as a silicone oxide film or a sillcon nitride film without contaminating the surface to be polished with an alkali metal such as sodlum ions and with no flaws, and the CMP abrasive is excellent in storage stability.

Description

Technical Field

5 [0001] The invention relates to a CMP abrasive usable in the production of semiconductor elements, a liquid additive for CMP abrasive and a method for polishing substrates. Particularly, it relates to a CMP abrasive usable in the step of planarizing the surface of a substrate, typically the steps of planarizing an interlayer insulating tim and forming shallow-trench separation, to an additive for the CMP abrasive and to a method for polishing substrates by using the CMP abrasive.

Background Art

[0002] In present ultra large scale Integrated circuits, packaging density is increasing, and various fine processing techniques have been studied and developed. Design rules have almady reached the order of sub half micron. Off (Chemical Mechanical Polishing) is one of the techniques developed to satisfy such a strict requirement for lineass. This techniques developed to satisfy such a strict requirement for lineass. This technique is essential for the production of semiconductor devices, typically for planarizing interlayer insulating films and for shallow-trench separation, because it can completely planarize layers to be exposed, reducing the burden on excousive techniques and stabilizing the production visid.

- 2003] Colloidal silica abrasives have been investigated as common CMP abrasives to be used in the production of semiconductor devices to planarize horganic insulating films, such as silican oxida insulating film, formed by planara-CVD (Chemical Vapor Deposition), tow pressure-CVD or the like. Colloidal silica abrasives may be produced by using silica particles, which are typically formed from tetrachicoreliane through thermal decomposition, and adjusting pH. Such abrasives, however, cannot polish in organic insulating films fast enough, and need higher polishing rate for their practical use.
- 28 [0004] In integrated circuita with design rules of 0.5 µm or more, devices were separated by LOCOS (Localized Oxidation of Silicon). As the processing measurements have become finer, shallow-trench separation has become used in response to the requirement for a technique giving narrower separation gap between devices. For shallow-trench separation, the europius parts of a silicon oxide film formed on a substrate are removed by CMP, and a stopper film reducing the polishing rate is provided under the silicon oxide film to stop polishing. The stopper film is typically amade of silicon intride, and the rates of polishing the silicon oxide film and the stopper film are preferably in a large ratio. Where conventional colloids silica ebrashws are used, the ratio between the rate of polishing the silicon oxide film and the stopper film is as small as the order of 3, and such abrasives cannot satisfy the requirements of practical shallow-tench separation.
- [0005] On the other hand, cerium oxide abrasives have been used for polishing photo masks or the surface of glass, is such as lenses. Having lower hardness as compared to silica particles and alumina particles, cerium oxide particles hardly make flaws on the polished surface and are sultable for finish mirror polishing. However, the cerium oxide abrasives for polishing glass surfaces cannot be used as abrasives for polishing semiconductors, because they contain a dispersant containing adulum salts.

40 Disclosure of Invention

[0006] An object of the invention is to provide a CMP abrasive, which can speedily polish a surface to be polished, such as a silicon oxide insulating film, without making flaws.

[0007] Another object of the invention is to provide a CMP abrasive, which can speedily polish a surface to be polsted, such as a silicon oxide insulating film, without contaminating the surface to be polished with aikall metals, such as sodium lone, nor making flaws.

[0008] Another object of the invention is to provide a CMP abrasive, which is more advantageous in that it can increase the ratio of the rate of polishing a silicon oxide insulating film to the rate of polishing a silicon nitride insulating film.

(9 [0099] Another object of the invention is to provide a CMP abrasive, which can speedily polish a surface to be polished, such as a sillcon oxide insulating film, without contaminating the surface to be polished with alkali metals, such as sodium lons nor making flaws and contains a cerium oxide slurry improved in storage stability.

(0010) Another object of the invention is to provide a CMP abreakw, which can speedity polish a surface to be polished, such as a sillicon oxide insulating film, without contaminating the surface to be polished with alkall metals, such as sodium lons, nor making flaws, and can increase the ratio of the rate of polishing a silicon oxide insulating film to the rate of polishing a silicon mixed insulating film to the rate of polishing as silicon tritical insulation film to 50 or more than the provided in the provided insulating film to the rate of polishing as silicon to the rate of polishing as silico

[0011] Another object of the Invention is to provide a liquid additive, which is to be used to give a CMP abrasive improved in storage stability.

- [0012] Another object of the invention is to provide a liquid additive for CMP abrasive to be used to improve the flatness of the polished surface of a substrate.
- [0013] Another object of the invention is to provide a method for polishing a substrate, which can polish a surface of the substrate without making flaws on its polished surface.
- [0014] Another object of the invention is to provide a method for polishing a substrate, which can speedily polish a surface to be polished, such as a silicon oxide insulating film, without making flaws, and can increase the ratio of the rate of polishing a silicon oxide insulating film to the rate of polishing a silicon nitride insulating film to 50 or more. Accordingly, the invention relates to:
- 10 (1) a CMP abrasive comprising
 - a cerium oxide siurry containing cerium oxide particles, a dispersant and water, and a liquid additive containing a dispersant and water:
- 15 (2) the CMP abrasive of (1), wherein each of the dispersants contained in the certum oxide stury and the liquid additive respectively is a polymer dispersant, which is a polymer containing ammonitum acrylate as a copolymerized introductor.
 - (3) the CMP abrasive of (1), wherein each of the dispersants contained in the cerium oxide sturry and the liquid additive respectively is a polymer dispersant, which is a polyarmonium-acrylate or a polyarmine-acrylate;
- (4) the CMP abrasive of (2) or (3), wherein the polymer dispersants have a weight average molecular weight of 100 to 50,000;
 - (5) the CMP abreative of (1), wherein the cerium oxide sturry contains 0.01 to 2.0 parts by weight of the dispersant relative to 100 parts by weight of the cerium oxide particles and contains 0.3 to 40 % by weight of the cerium oxide particles based on the cerium oxide sign.
- 25 (6) the CMP abrasive of any one of (1) to (5), wherein the cerium oxide slurry is pH 6 to 10;
 - (7) the CMP abrasive of any one of (1) to (6), which is 50 or more in ratio of rate of polishing a silicon oxide film to rate of polishing a silicon nitride film:
 - (8) a liquid additive for CMP abrasive comprising a dispersant and water,
 - (9) the liquid additive for CMP abrasive of (8), which contains 1 to 10 % by weight of the dispersant;
- (10) the liquid additive for CMP abrasive of (9), wherein the dispersant is a polyammonium-acrylate or a polyamineacrylate;
 - (11) the liquid additive for CMP abrasive of (9), wherein the polyammonium-acrylate or the polyamine-acrylate has a weight average molecular weight of 1,000 to 100,000:
- (12) the liquid additive for CMP abrasive of (11), wherein the polyammonium-ecrylate or the polyamine-acrylate has a molecular weight distribution (weight average molecular weight/number average molecular weight) of 1,005 to 1,300;
 - (13) the liquid additive for CMP abrasive of (12), wherein the polyammonium-acrylate or the polyamine-acrylate contains 10 mol % or less of free ammonia or a free amine, which does not form a salt;
 - (14) the liquid additive for CMP abrasive of (10), which is pH 4 to 8;
- 40 (15) the liquid additive for CMP abrasive of (10), which has a viscosity of 1.20 to 2.50 mPa s;
 - (16) a method for polishing a substrate, comprising holding a substrate having, formed thereon, a film to be polished against a polishing pad of a polishing platen, followed by pressing, and moving the substrate and the polishing platen while supplying the CMP abrasive of any one of (1) to (7) in between the film to be polished and the polishing pad to thereby polish the film to be polished; and
- (17) the method of (16), wherein the substrate to be polished has at least a silicon oxide film or a silicon nitride film formed thereon.

Best Mode for Carrying Out the Invention

- 50 [0015] Cerium oxide may be produced by the oxidation of a cerium compound, such as carbonate, nitrate, sulfate or oxalate of cerium. Conventional cerium oxide abrasives for polishing silicon oxide films formed by TEOS-CVD or the like contain monocrystatiline cerium oxide particles with large primary particle sizes, and tend to make polishing flaws. Therefore, the cerium oxide particles to be used in the Invention are not limited in the method of production, but are preferably polycrystals that are aggregates of monocrystals of 5 mm to 300 nm. For polishing semiconductor chips, the cerium oxide particles preferably contain as little as 10 ppm or less of aliest imetals and halogens.
 - [0016] The methods usable in the invention to produce the certum oxide powder include burning or oxidation using hydrogen peroxide or the like. Preferred burning temperatures range from 350 to 900°C. The raw material suitable for the method is certum carbonate.

[0017] The above method gives aggragates of certum oxide particles, which are then preferably pulverized mechanically. Examples of preferrad pulverizing methods include dry grinding using a jet mill, and wet grinding using a planetary bead mill.

[0018] The cerium oxide slurry to be used in the invention is obtainable, for example, by dispersing a composition comprising the cerium oxide particles having the above characteristics, a disparsant for dispersing the cerium oxide particles in water and water. The content of the cerium oxide particles is not limited but is preferably 0.3 to 40 % pwelght, more preferably 0.5 to 20 % by weight to handle the dispersion easily, in the CMP abrastve obtainable by my will be certum oxide surroy and a liquid additive, the content of the cerium oxide particles is preferably 0.01 to 10 % by weight, more preferably 0.10 to 5 % by weight.

[0019] The dispersant to be used in the Invention comprises one or two compounds selected from polymer dispersants, water-soluble anionic surfactants, water-soluble nonlonic surfactants, water-soluble cationic surfactants and water-soluble amphoteric surfactants. For polishing semiconductor chips, the dispersant preferably contains as little gas 10 ppm or less of alkali metals, such as sodium ions and potassium ions, hadogens and suiting.

[0020] Examples of polymer dispersants include polymers of unsaturated carboxylic acids, such as acrylic acid, methacylic acid or mailed acid or armined such or amine salts of the polymers: copolymers of an unsaturated carboxylc acid, such as acrylic acid, methacrylic acid or mailed acid, with a copolymertzable monomer, for example, an alkyl acrylate, such as methyl acrylate or ethyl acrylate, a hydroxylatyla acrylate, such as methyl acrylate or ethyl acrylate, and alkyl methacrylate, alkyloxylate, which are hydroxylatily methacrylate, acid not set in the polymers or copolymers may be converted into ammonium salts or the copolymers. The unsaturated acroxylate acid moleties in the polymers or copolymers may be converted into ammonium salts either before or after polymerization. The polymers and copolymers preferably contain 1 to 100 mol%, more preferably 10 to 100 mol%

[0021] Preferred dispersants are polymers containing ammonium acrylate as a copolymerized ingredient, polyarmonium-acrylates and polyarmine-acrylates. Polyarmonium-acrylates and polyarmine-acrylates preferably have weight average molecular weights of 1,000 to 10,000, more preferably 3,000 to 80,000, turber preferably 10,000 to 40,000. If the weight average molecular weight is less than 1,000, cerium oxide particles may aggregate, and if more than 1,000, polishing rate ratio may be reduced. The polyarmonium-acrylates and polyarmine-acrylates preferably have molecular weight distributions (weight average molecular weight withumber average molecular weight of 1,005 to 3,000, more preferably 1,100 to 1,250, further preferably 1,150 to 1,200. If the molecular weight distribution is less than 1,005, the cerium oxide particles may aggregate, and if more than 1,300, the pollaring rate ratio may be reduced. Herein, the weight average molecular weight and number average molecular weight are measured through get permeation other.

matography, based on the calibration curve of a standard, polystyrene.

[0022] The polyammonium-acrylates and polyamine-acrylates are obtainable through neutralization of a mixture comprising a polyacopito acid and an equimolar amount of ammonia or an amine relative to carboxyl groups, and, in view of high flatness, preferably contain as fittle as 10 mol% or less of fine ammonia or amine forming no satis (that is, at least 90 mol% of the carboxyl groups of polyacrylic acid are preferably neutralizad). The amount of the fire ammonia or amine in the fittle acid and the polymer and quantitatively determining the amount of ammonia or amine in the fittrate.

[0023] Examples of water-soluble anionic surfactants include triethanolamine lauryl sulfate, ammonium lauryl sulfate and triethanolamine polyoxyethylene alkyl ether sulfates.

[0024] Examples of water-soluble nonlonic surfactants include polyoxyethylene lauryl ether, polyoxyethylene cetyl ether, polyoxyethylene stearly ether, polyoxyethylene oleyl ether, polyoxyethylene higher alcohol ethers, polyoxyethylene octyl phemyl ether, polyoxyethylene ender phemyl ether, polyoxyethylene ender phemyl ether, polyoxyethylene edher, polyoxyethylene edher, polyoxyethylene edher and ender ether edher edher ether, polyoxyethylenesolibitan monostear-tae, ploxyethylenesolibitan trialseartae, polyoxyethylenesolibitan trialseartae, polyoxyethylenesolibitae, polyoxyethyleneso

(2025) Examples of water-soluble amphoteric surfactants include laurylbetaine, stearylbetaine, lauryldmethylamine oxide and 2-alkyl-N-carboxymethyl-N-hydroxyethylmidazolinlumbetaine.

[0026] In view of the dispersibility of particles in the certum oxide alurry, the prevention of their sedimentation and the relationship between polishing flaws and the amount of the dispersants added, the amount of the dispersant added in the certum oxide surrive preferably 0.01 to 2.0 parts by weight relative to 100 parts by weight of certum oxide particles. [0027] Among the dispersants described above, the polymer dispersants preferably have weight average molecular weights of 100 to 100,000, more preferably 100 to 50,000, further preferably 10,001 to 10,000, as measured through get permeation chromatography based on the calibration curve of the standard, polystyrene. If the molecular weight of the dispersant is too low, silicon oxide films or silicon nitride films may not be polished speedily enough, and if it is

too high, the cerium oxide slurry may be too viscous and lose storage stability.

[0028] The pH of the cerium oxide airury is preferably 5 to 10. If the pH is too low, a liquid mixture of the cerium oxide airury and a liquid additive may lose storage stability and make polishing flaws on polished silicon oxide or silicon florid films, and if the pH is too high, a liquid mixture of the cerium oxide sturry and a liquid additive may also lose storage stability and make polishing flaws on polished silicon oxide or silicon nitride films. The pH may be adjusted by adding an aque ous ammonia and stirring.

[0029] The cenum oxide particles can be dispersed in water by using a common stirrer or others, such as a homogenizer, an ultrasonic disperser or a wet ball mill.

[0030] The cerium oxide particles in the slurry thus produced preferably have an average particle size of 0.01 to 1.0 µm. This is because an abrasive with a too small average particle size has a considerably low polishing rate, and that with a too iscree everage particle size tends to make flaws on the polished film.

10031] The liquid additive for CMP abrasive of the invention comprises a dispersant and water. The dispersant is for dispersing the cerium oxide particles contained in the above-described cerium oxide particles possible or the polishing rate ratio and the high flatness of the polished surface, the dispersants suitable for the cerium oxide surry in water. In view of the polished is unlikely to the dispersants suitable for the cerium oxide surry are also suitable for the liquid additive. The dispersants used in the cerium oxide surry and the liquid additive may be identical with or different from each other. The concentration of the dispersant in the liquid additive is preferrable to 10.5% by weight. If it is less than 1 % by weight, the polished surface may be less flat, and if more than 10 % by weight, the polished surface may be less flat, and if more than 10 % by weight,

[0032] The CMP abrasive of the invention is used so that the certum oxide sturry and the liquid additive prepared apart from each other are mixed at the time of polishing. If the certum oxide sturry and the liquid additive are stored in a form of a mature, the certum oxide particles will aggregate, thereby making polishing flaws and causing a change in polishing rate. Therefore, the liquid additive and the certum oxide sturry are supplied on a polishing platen separately and mixed thereon, or mixed immediately before polishing and then supplied onto a polishing platen. The mixing tole between the certum oxide sturry and the liquid additive is not particularly limited so far as the desired concentrations

[033] In view of the dispersibility of particles in the slurry, the prevention of their sedimentation and the relationship between polishing flaws and the amount of the dispersant added, the amount of the dispersant in the liquid addition relative to the cordium oxide is preferably 0.01 to 2000 parts by weight, more preferably 0.01 to 1000 parts by weight, further preferably 0.01 to 500 parts by weight, relative to 100 parts by weight of the cerium oxide particles in the cerium oxide slurry.

10034] The specific gravity of the liquid actifive is preferably 1.005 to 1.050, more preferably 1.007 to 1.040, further preferably 4.007 to 1.040 further preferably 4.007 to 1.040 further preferably 4.007 to 1.040 further preferably 5.007 for 1.040 further preferably 4.007 to 1.040 further preferably 4.007 to 1.040 further preferably 6.007 fur

[0036] The inorganic insulating film for which the CMP abrashe of the invention is used may be formed, for example, by low pressure CVD, or plasma CVD. To form a silicon oxide film by low pressure CVD, monosiliane: SiH_a is used as an Si-source, and oxygen: O₂ as an oxygen-source, and SiH_a-O₂ exidation is carried out at a low temperature of 400°C or lower. After CVD, heat treatment at a temperature not higher than 1000°C may optionally be carried out. Where phosphorus: P is doped to planarize the surface by high temperature reflow, an SiH_a-O₂-PH_a reaction gas is preferably used. Plasma CVD is advantageous in that chemical reactions requiring high temperatures under normal, thermal equilibrium can undergo at lower temperatures. The methods for generating plasma include two types of capacitive coupling and inductive coupling. Examples of reaction gases include an SiH_a-N₂-O gas comprising SiH_a as an Si-source and N₂O as an oxygen-source; and a TEOS-O₂ gas containing tetraethoxyellane (TEOS) as an Si-source (TEOS-plasma CVD). The preferred temperature of the substrate ranges from 250 to 400°C, and preferred reaction pressure ranges from 67 to 400 Pa. As described above, the silicon oxide film to be used in the invention may be doped with other elements, such as phosphorus or boron.

[0037] To form a silicon nitride film by low pressure CVD, dichlorosilane: SIH₂Ci₂ is used as an Si-source, and ammonia: NiH₃ as a nitrogen-source, and the SiH₂Ci₂NiH₃ outdation is carried out at a high temperature of 900°C. An example of the reaction gas for plasma CVD is an SiH₄NiH₃ gas comprising SiH₄ as an Si-source and NiH₃ as a

nitrogen-source. The preferred temperature of the substrate ranges from 300 to 400°C.

[0038] The substrate to be used may be a semiconductor substrate bearing circuit devices and wiring patterns formed thereon, or a semiconductor substrate which bears circuit devices formed thereon and is further coated with a silicon oxide film layer or a silicon nitride film layer. Polishing a silicon oxide film layer are allicon nitride film layer through oxide film layer or a silicon nitride film layer through oxide film layer through the surface of the sellicon oxide film layer, to planarize whole the surface of the selliconductor substrate, it is also applicable for shallow-trench separation. Shallow-trench separation. Shallow-trench separation needs a ratio of the rate of polishing a silicon notified film polishing rate of 10 or more. If the ratio is too small, the difference between the silicon oxide film-polishing rate and the silicon intride film-polishing rate will be too small, to stop polishing at a position predetermined for shallow-trench separation. If the ratio is 50 or more, polishing can be stopped easily by the further reduced polishing rate of silicon nitride film, and the CMP abrasive with such a ratio is room suitable for shallow-trench separation.

[0039] The polishing apparatus to be used may be a common one, which has a holder for holding a semiconductor substrate and a platen (equipped with a motor or the like capable of changing rotational speed) applied with a polishing pad. The material of the polishing pad may be any one, such as a nonwoven fabric, a polyurethane foam or a porous fluorine resin. The polishing pad is preferably grooved to collect the CMP abrasive in the grooves. The polishing conditions are not limited, but the rotational speed of the platen is preferably as low as 200 rpm or less to prevent the semiconductor substrate from being emitted. The pressure applied to the semiconductor substrate is preferably 1 kg/ cm² or less not to make polishing flaws. For shallow-trench separation, polishing should make few flaws. During pol-Ishing, the slurry is continuously supplied to the polishing pad by some means, such as a pump. Not limitative but preferred amount of the slurry supplied is such that the surface of the polishing pad is always coated with the slurry. [0040] After pollshing, the semiconductor substrate is preferably washed well in running water and then dried after blowing away the water droplets from the semiconductor substrate by a spin dner or the like. Thus a planarized shallowtrench structure is formed. Subsequently, aluminum wiring is formed on the silicon oxide insulating film layer, and a silicon oxide insulating film is again formed between and on the wiring by the same method as described above and polished by using the CMP abrasive to smooth out the unevenness on the insulating film surface, thereby planarizing whole the surface of the semiconductor substrate. The process is repeated to produce a semiconductor with desired lavers.

[0041] The CMP abrasive of the invention can polish not only the silicon oxide film formed on a semiconductor substrate but also an inorganic insulating film formed on a wiring board bearing a predetermined writing, such as a silicon oxide film, glass or silicon intride; optical glass, such as photo masks, lensee and prisme; inorganic condeautor films, such as ITC, optical integrated circuits, optical switching devices and optical guides, which are made of plass and orystalline materials; the end faces of optical fibers; optical monocystals, such as softilitators; solid-state armonocystals; sapphire substrates for blue laser LED; semiconductor monocrystals, such as SIC, GaP and GaAS; glass substrates for magnetic class: and magnetic heads.

[0042] Hereinafter, the Invention will be described in more detail referring to Examples and Comparative Examples, which however do not limit the scope of the invention.

Preparation 1 (Preparation of cerium oxide particles)

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[0043] 2 kg of certum carbonate hydrate was placed in a platinum vessel and burned in the air at 700°C for 2 hours, to give about 1 kg of yeallowish white powder. The powder was identified to be certum oxide by X-ray diffractionetry. The certum oxide powder was mixed with delonized water to 10 % by weight concentration, and pulverized with a horizontal wet ultrafine dispersing-pulverizer at 1400 pm for 120 minutes. The resulting flouid abreative was heated to 110°C for 3 hours to give dry certum oxide particles. The certum oxide particles were polycrystale comprising 10 60-cm-particle size primary particles as observed by a transmission electron microscope, and had a specific surface area of 35.6 m/g as measured by the BET method.

Preparation 2 (Preparation of cerium oxide particles)

[0044] 2 kg of certum carbonete hydrate was placed in an platinum vessel and burned in the air at 700°C. for 2 hours, to give about 1 kg of yellowish white powder. The powder was identified to be certum oxide by X-ray diffractiometry. 1 kg of the certum oxide powder was dry-ground with a jet mil. The certum oxide perticles were polycrystate comprise of 10 miles of the certum oxide powder was dry-ground with a jet mil. The certum oxide perticles were polycrystate comprise of 10 miles of 10 mile

Preparation 3 (Preparation of certum oxide slury)

[0045] 125 g of the cerium oxide particles prepared in Preparation 1, 3 g of a 40-wt% aqueous solution of an ammonium satif of polyacytic acid copolymen, which was a 3-1 -copolymentzation product of acrylic acid and methyl acrylication and had an weight average molecular weight of 10,000, and 2372 g of delonized water were mixed, and ultrasonized, dispersed with attiring. The dispersing was conducted for 10 minutes with an ultrasonic frequency of 40 kHz. The resulting situry was filtered through a 0.8-µm filter, and delonized water was added thereto to give a 2-wt% certimoxide sutrry (A-1). The pH of the certimoxide situry (A-1) was 8.5. The certimoxide situry (A-1) contained particle with an average particle size of as small as 0.20 µm as determined from their particle size distribution measuring apparatus. 9.5.0 % of the particles were 1.0 µm or less.

Preparation 4 (Preparation of cerium oxide slurry)

[0046] A cerium oxide siurry (A-2) was prepared in the same manner as in Preparation 3, except the cerium oxide particles prepared in Preparation 1 were replaced by the cerium oxide sturry prepared in Preparation 2. The pH of the cerium oxide sturry (A-2) except solury (A-2) except sol

Preparation 5 (Preparation of cerium oxide particles)

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[0047] 2 kg of cerlum carbonate hydrate was placed in a platinum vessel and burned in the air at 900°C for 2 hours, to give about 1 kg of yellowish white powder. The powder was identified to be cerlum oxide by X-ray diffractionetry. It kg of the cerlum oxide powder was dry-ground with a jet mill. The cerlum oxide perdices were monocrystate of to 150 nm in particle size as observed by a transmission electron microscope, and had a specific surface area of 18.5 m/sg as measured by the BET method.

Preparation 6 (Preparation of cerium oxide slurry)

[0048] A cerium oxide siurry (8-1) was prepared in the same manner as in Preparation 1, except the cerium oxide particles prepared in Preparation 1 were replaced by the cerium oxide particles prepared in Comparative Preparation 1. The pith of the cerium oxide siurry (6-1) was 8-4. The cerium oxide siurry (6-1) oxontained particles with an average particle size of as small as 0.35 µm as determined from their particle size of distribution. 85.5 % of the particles were 1.0 µm or less.

35 Examples 1-10 and Comparative Examples 1 and 2

[0049] Cerlum oxide sturries and liquid additives were prepared to prepare the CMP abrasives as shown in Table 1, and mixtures of a cerlum oxide sturries and a liquid additive were used for polishing an insulating film in the manner described below. The results are listed in Table 1.

[0050] The liquid additive used in each of Examples 1-5, 7 and 9 was prepared by dissolving in deionized water a predetermined amount of the same dispersant as that used in the cerium oxide slurry of Example 1.

[0051] The dispersant used in Examples 6, 8 and 10 was a polyammonium-acrylate having a weight average molecular weight of 10,000, a number average molecular weight of 8,333, a molecular weight distribution of 1.2 and a content of free ammonium of 4.3 mol%. The liquid additive used in Example 6 had a viscosity of 1.46 mPa-s and a specific gravity of 1.010.

[0052] In Comparative Example 2, the same cerium oxide slurry and liquid additive as those used in Example 1 were previously mixed, and the mixture was used for polishing an insulating film one day after.

(Polishing of Insulating film)

[0053] A 125-mm-diameter sillcon wafer with a silicon oxide film formed thereon by TEOS-plasma CVD was fixed to a holder to which an attraction pad for fixing substrates was bonded, and was then set, with the surface of the insulating film directed downwardly, on a platen to which a polishing pad made of a porous urefulene resin was bonded. A weight was then placed on it to produce a load of 300 g/cm². The insulating film was polished by rotating the platen at 40 pm/for 2 minutes while rededing a certim oxide sturry (solid content: 25 to Weight) and a liquid additive separately both at a rate of 25 m/min and dropping them as one liquid onto the platen by controlling nozzies so that they joined together just above the platen. After polishing, the wafer was removed from the holder, washed with running water well, and then with an ultrasonic cleaner for 20 minutes. After washing, water droplets were removed by a spin diser,

and the wafer was dried for 10 minutes in a 120°C drier. The change in the film thickness before and after polishing was measured with a photo-interierent film thickness measuring apparatus to determine the polishing rate, [0054] In place of the silicon oxide film formed by TEOS-plasma CVD, a silicon nitride film produced by low pressure

CVD was polished in the same manner under the same conditions, and the change in the film thickness before and after polishing was measured, to determine the polishing rate. The results of the measurements of film thickness show that the silicon oxide film produced by TEOS-plasma CVD and the silicon nitride film produced by low pressure CVD were made uniform in thickness all over the waters. No flaws were observed on the surfaces of the insulating films by visual observation under a mercury-vapor lamp, but the surfaces were further observed precisely with an apparatus for examining the appearance of wafers (trade name: OLYMPUS Al-2000, produced by Olympus Optical Co., Ltd.), [0055] Similarly, a sillcon oxide film having 20-µm-square 5,000-Å-high projections at 100-µm distances was polished, and the degree of dishing was measured at intermediate points between the polished projections to evaluate

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Table 1-1		Example 6	A		+	8.5 Acrylic acid / methyl acrylate		am Trace	10,000			3	8.9		1,800		20		06		3		60.0	80				
		Example 5 (A-2) 10-60 Polyczystal			8.7 Acryllo acid / methyl acrylate		= 3/1	000'01 .				2	7.5		2,000		50		100		0.05		20.10	130				
		(A-2) (A-2) 10-60 Polyczystal		7.8	Acres 4 c. c. d.	/ methyl	acrylate	= 3/1		10,000			2	7.5	2,000			20		100			0.05		130			
	Fysum 1a 3	(1,8,1)	(7-8)	10-60 Polycrystal 8.5		Acrylic acid / methyl acrylate = 3/1		10,000		-	7.7		1,500			20		75		0.05		100						
	Example 2	(A-1)	(A-1) 10-60 Polycrystal 8.5		Acrylic acid / methyl acrylate = 3/1		= 3/1	10,000		-	2		2,000			50		100			0.05	120						
	Example 1	(A-1)		10-60	rotyonystal	8.5	Acrylic acid	/ methyl	acrylate	= 3/1		10,000		-	7.3	?!	, 000	9		\$			20		100	6.02	150	
		Name	Secondary	particle size	(mm)	PH.		Dispersant		Motoh	average	molecular	weight	Conc. (wt%)	Ha	Plasma-CVD-TEOS-s111con	oxide film polishing	n)	re-CVD-	silicon nitride film	polishing rate (A/min)	Polishing rate ratio	xdde film /	tride film)	Flaws on polished oxide	er/cm²)	Degree of dishing (A)	
		Cerlum oxide slurry (500 g)					Liquid additive (500 g)									Plasma-CVD	oxide film	rate (A/min	Low pressure-CVD-	silicon ni	polishing	Polishing	(silicon oxide film / silicon nitride film)		Flaws on p	film (number/cm ²)	Degree of	

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[0056] As apparent from Table 1, the CMP abrasive and method for polishing substrates according to the invention can polish a surface to be polished, such as a silicon oxide film or a silicon nitride film, without contaminating the surface to be polished with alkall metals, such as sodium ions, nor making flaws, and further can increase ratio of (silicon oxide film polishing rate)/(silicon nitride film polishing rate) to 50 or matarets.

Industrial Applicability

[0057] The CMP abrasive of the invention is sultable for polishing methods used in the production of semiconductor elements, particularly for polishing exbetrates for shallow-tranch separation because it can speedily polish a surface to be polished, such as a silicon oxide tilm, without making flaws.

[0088] The CMP abrasive of the invention is as well advantageous in that it does not contaminate the surface to be polished with alkali metals, such as sodium ions, and can increase the ratio of (silicon oxide film polishing rate).(silicon nitride film polishing rate).

[0059] The CMP abrasive of the invention is suitable for polishing methods used in the production of semiconductor selements because it can improve the storage stability of cerium oxide siurnies.

[0080] The method for polishing substrates of the invention is suitably applicable in the production of semiconductor elements because it excels in polishing speedily a surface to be polished, such as allicon oxide film, without making flaws.

Cialms

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- 1. A CMP abrasive comprising
 - a cerium oxide siurry containing cerium oxide particles, a dispersant and water; and a liquid additive containing a dispersant and water.
- 2. The CMP abrasive of claim 1, wherein each of the dispersants contained in the certum oxide siumy and the liquid additive respectively is a polymer dispersant, which is a polymer containing ammonium acrylate as a copolymerized ingredient.
 - The CMP abrasive of claim 1, wherein each of the dispersants contained in the certum oxide siurry and the liquid additive respectively is a polymer dispersant, which is a polyammonium-acrylate or a polyamine-acrylate.
- The CMP abrasive of claim 2 or 3, wherein the polymer dispersants have a weight average molecular weight of 100 to 50,000.
 - The abrasive of claim 1, wherein the cerium oxide sturry contains 0.01 to 2.0 parts by weight of the dispersant relative to 100 parts by weight of the cerium oxide particles and contains 0.3 to 40 % by weight of the cerium oxide particles based on the cerium oxide sturry.
 - 6. The CMP abrasive of any one of claims 1 to 5, wherein the cerlum oxide slurry is pH 8 to 10.
- The CMP abrasive of any one of claims 1 to 6, which is 50 or more in ratio of rate of polishing a silicon oxide film to rate of polishing a silicon nitride film.
 - 8. A liquid additive for CMP abrasive comprising a dispersant and water.
 - 9. The liquid additive for CMP abrasive of claim 8, which contains 1 to 10 % by weight of the dispersant.
 - The liquid additive for CMP abrasive of claim 9, wherein the dispersant is a polyammonium-acrylate or a polyamineacrylate.
- 11. The liquid additive for CMP abrasive of claim 9, wherein the polyammonium-acrylate or the polyamine-acrylate has a weight average molecular weight of 1,000 to 100,000.
 - 12. The liquid additive for CMP abrasive of claim 11, wherein the polyammenium-acrylate or the polyamine-acrylate has a molecular weight distribution (weight average molecular weight/number average molecular weight) of 1.005

to 1.300.

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- 13. The liquid additive for CMP abrasive of claim 10, wherein the polyammonium-acrylate or the polyamine-acrylate contains 10 mol % or less of free ammonia or a free amine, which does not form a sait.
- 14. The liquid additive for CMP abrasive of claim 10, which is pH 4 to 8.
- 15. The liquid additive for CMP abrasive of claim 10, which has a viscosity of 1.20 to 2.50 mPa s.
- 9 16. A method for polishing a substrate, comprising holding a substrate having, formed thereon, a film to be polished against a polishing part of a polishing puriace platen, followed by pressing, and moving the substrate and the polishing platen while supplying the CMP abrasive of any one of claims 1 to 7 in between the film to be polished and the polishing pad to thereby polish the film to be polished.
- 15 17. The method of claim 16, wherein the substrate to be polished has at least a silicon oxide film or a silicon nitride film formed thereon.